DEPARTMENT OF HEALTH AND HUMAN SERVICES

PUBLIC HEALTH SERVICE

APPLICATION FOR A VARIANCE FROM 21 CFR 1040.11(c) FOR A LASER LIGHT SHOW, DISPLAY, OR DEVICE

Form Approved: OMB No. 0910-0025. Expiration Date: October 31, 1991. See Page 4 for OMB Statement.

. FOOD AND DRUG ADMINISTRATION			SER LIGHT SHOW, D OR DEVICE	ISPLAY,	DOCKET NUMBER				
NOTE: No laser light show, projection system, or devapplication in accordance with 21 CFR 1010.4	vice may vary f l.	rom comp	oliance with 21 CFR 1040).11(c) in design o	r use without the approval of this				
1. Check all applicable boxes and type or print the requested information. 2. Submit an original and four (4) copies.		INSTRU 3. Ma an 4. En	CTIONS ail your application कि ti d Drug Administration, ter docket number if as	ne Dockets Mana Room 4-62, 5600 signed	gemen Bradch (HFA-305), Food Fishers Lane, Rockville, MD 20857.				
1. NAME OF COMPANY ACCESS ELECTRODICS									
2. ADDRESS OF COMPANY (Include ZIP CODE) (If P.C). Box is used, ii	nclude act	tual street address also.)					
3900 NORENE LN LOUISVILLE KY 40219									
3. NAME AND TITLE OF RESPONSIBLE PERSON	4. TELEP		. (Include area code)		5. DATE OF SUBMISSION				
ALLEN OWENS			1920		70-10-99				
6. THE APPLICANT REQUESTS THE VARIANCE TO BE IN EFFECT FOR A PERIOD OF									
7.			IPTION AND USE						
a. LIST NAME AND/OR MODEL NUMBER(S) FOR THE					•				
Η Μερίοω 60 Χ Αρίου, Μείος Gr b. PRODUCT FOR WHICH A VARIANCE IS REQUESTED □ A laser display device □ A projector for a laser light show ☑ A laser light show □ Other (Specify)	f. PRÓDUCT IS INTENDED TO BE USED AT ANY ONE LOCATION ☐ More than 15 days ☐ More than 5 but not more than 15 days ☑ Less than 5 days								
c. Projectors are intended for sale, lease, or loan to other laser light show producers			g. TOUR IS INTENDED TO RUN FOR More than 6 months 1-6 months Less than one month Not applicable (Not a tour) Other (Specify) h. PRODUCT UTILIZES THE FOLLOWING LASER EFFECTS Front screen projections Rear screen projections Multiple reflection/diffraction effects Audience scanning (Also includes scanning any accessible uncontrolled areas) Reflections from stationary mirrors or mirrored surfaces (Beam Matrices) Stationary irradiation of rotating mirror balls, etc. Scanning irradiation of rotating mirror balls, etc.						
d. PRODUCT IS INTENDED FOR USE IN A Planetarium or other dome projection structure Theater Hotel/motel ballroom or meeting room Store displays Trade show or convention Discotheque or night club Pavilion Indoor arena Outdoor arena Museum Outdoor unenclosed area Mother (Specify) ALL OF The Above									
☐ At only one (Fixed) location At a variety of (Tour) locations ☐ Other (Specify)			Fog, smoke, or other scattering enhancement effects Other (Specify)						
8.	LAS	ER RADIA	TION LEVELS						
LASER MEDIUM (Ar, He-Ne, etc.)	WA	AVE LENG	THS (nm)		PEAK POWER (watts)				
AMERICAN 60x ARLON	496,5 70 5		514,5 AM	250n	w				
Melles GRIOT HELIUM NEON 632.80					ω				
9. IF ANY LASER RADIATION IS PULSED OR SCANNED), GIVE THE PU	LSE DURA	TION AND RATE AND S	CANNING FREQU	ENCY AND AMPLITUDE				
10. REASON FOR REQUESTING VARIANCE									
Compliance with the limits of 21 CFR 1040.1 limit the output power to the extent that th Other or additional explanation (Specify)	1(c) would rest e desired effe	trict the in cts would	itended use of the prod not be sufficiently visibl	uct because comp e	oliance would				

11.	MANNER IN WHICH IT IS PROPOSED TO DEVIATE FROM THE REQUIREMENTS OF THE APPLICABLE STANDARD								
	ē	It is proposed to deviate from the provisions of 21 CFR 1040.11(c) in that the accessible emission level would exceed the accessible emission limits specified in 21 CFR 1040.11(c).							
	[It is proposed to deviate from the provision of 21 CFR 1040.11(c) as follows:						
12.	AD	V	ANTAGES TO BE DERIVED FROM SUCH DEVIATION						
	[X	Laser light shows and displays are accepted popular media in entertainment and the arts. Use of power levels in excess of the limits imposed by 21 CFR 1040.11(c) is necessary to achieve the required effects in these media.						
	[J	Other or additional advantages (describe and explain).						
13.	EXI	PL.	AIN THE ALTERNATE MEANS OF RADIATION PROTECTION TO BE PROVIDED. (Check as many boxes as apply. In Item 14 "Remarks," justify ar s not checked, using additional sheets as necessary. State any other means of radiation protection that will be used.)						
	a. [M	All laser products, systems, shows, and projectors will be certified to comply with 21 CFR 1040.10 and the conditions of this variance and will be reported as required by 21 CFR 1002.10 and 1002.12 using the reporting guides provided for such purpose. These actions will be accomplished prior to any introduction into commerce.						
	b. (Ø	Effects not specifically indicated in this variance application will not be performed. No other effects will be added until an amendment to the variance has been obtained and the required reports or supplements, as applicable, have been submitted.						
	c . [æ	Scanning, projection, or reflection of laser and collateral radiation (<i>Light show radiation</i>) into audience or other accessible uncontrolled areas will not be permitted except for diffuse reflections produced by the atmosphere, added atmospheric scattering media, and target screens.						
	d. [DX.	Laser radiation levels in excess of the limits of Class I will not be permitted at any point less than 3.0 meters above any surface upon which persons other than operators, performers, or employees are permitted to stand or 2.5 meters below or in lateral separation from any place where such persons are permitted to be. Operators, performers, and employees will not be required or allowed to view radiation above the limits of Class I or be exposed to radiation above the limits specified in 21 CFR 1040.11(c).						
	e. [K	Any product which relies on scanning to meet access, exposure, or product class limits will incorporate a scanning safeguard system which directly senses scanner motion and which will react fast enough to preclude exceeding the applicable limit.						
	f. (M	All laser light shows shall be under the direct and personal control of trained, competent operator(s). The operator(s) will:						
			(1) Immediately terminate the emission of light show radiation in the event of any unsafe condition;						
			(2) Be located where all beam paths can be directly observed at all times; and						
			(3) Be an employee of the variance holder who will be responsible for the training and the conduct of the operator.						
	g.	M	The maximum laser projector output power will not exceed the level required to obtain the intended effects.						
	h.	Ø	The projection system (i.e., the projector and all other components used to produce the lighting effects) will be securely mounted or immobilized to prevent unintended movement or misalignment. Beam masking will be provided as an inherent part of the system design to prevent overfilling of screens, beam stops, targets, etc.						
	i.		Laser projectors will not be delivered to any other party under an agreement of sale, lease, or loan unless and until the recipient demonstrates that they have a variance in effect at the time of delivery that permits them to produce laser light shows incorporating such projector(s).						
	j.	Ø	In addition to the requirements of 21 CFR 1040.10(h), the manufacturer of laser projectors/systems will provide to parties who purchase, lease, or borrow the equipment, adequate users' instructions for safe installation and operation which explain the responsibility of the recipient as an independent light show manufacturer to submit the required reports and apply for and obtain a variance from CDRH prior to introduction into commerce of any laser light shows.						
	k.	2	The requirements of 21 CFR 1002.30(a)(1) and (2) will be accomplished through the use of written procedures for setup, alignment, testing, and performance of each show. These procedures will be in sufficient detail to ensure compliance with 21 CFR 1040.10, the conditions of the variance, and the control of access to radiation areas using the procedures described in the ANSI Z136.1 standard for the safe use of lasers (American National Standards Institute, 1430 Broadway, New York, NY 10018) or any other equivalent user consensus standard and, where applicable, state or local requirements. Laser radiation areas which can contain radiation levels above the limits specified in 21 CFR 1040.11(c) will be clearly identified by the posting of warning signs and/or restricting access through physical means (such as pressure switches, photo cells, barriers, guards, etc.). These requirements apply to temporary areas (such as during set up and alignment procedures) and to final or permanent areas. The variance holder will retain the records of these procedures and the results of all tests as required by 21 CFR 1002.31. A copy of the variance application, the approval letter, current procedures, and records relating to each particular show will be with the operator or other responsible individual and will be made available for inspection by FDA and other responsible authorities.						

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- Advance written notification will be made as early as possible to appropriate federal, state, and local authorities providing show itinerary with dates and locations clearly and completely identified, and a basic description of the proposed effects including a statement of the maximum power output intended. Such notifications will be made, but not necessarily be limited, to:
 - (1) The Center for Devices and Radiological Health, Office of Compliance and Surveillance (HFZ-312), 1390 Piccard Drive, Rockville, MD 20850, providing the initial and closing dates for fixed installations and the itinerary for mobile shows. In addition, unless all aspects of each show have been reported and accession numbers clearly referenced, each notice will include detailed descriptions of each show and a listing of all effects to be performed in sufficient detail to confirm compliance with the regulations and this variance.
 - (2) The Federal Aviation Administration (FAA) for any projections into open airspace at any time (i.e., including set up, alignment, rehearsals, performances, etc.). If the FAA objects to any laser effects, the objections will be resolved and any conditions requested by FAA will be adhered to. If these conditions cannot be met, the objectionable effects will be deleted from the show.
 - (3) State and local radiation control offices/agencies for all shows to be performed within their jurisdictions. All requirements of state and local law will be satisfied and any objections raised by local authorities will be resolved or the effects deleted. (A list of federal and state offices is available from the Center for Devices and Radiological Health upon request.)

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CERTIFICATION

I CERTIFY that all of the above information and statements are true, complete, and correct to the best of my knowledge and acknowledge that my variance application may be denied or my variance may be revoked if this application is found to be false, misleading, or incorrect in any material way. I have submitted and will submit all reports required by 21 CFR 1002.10 and 1002.12 on the laser equipment and show(s). I further understand that I may be required by regulation or by the Director, Center for Devices and Radiological Health, to supply such other information as may be necessary to evaluate and act on this application.

15. SIGNATURE

Aller Overst

16. NAME (Type or Print)

ALLW OWENS

17. TITLE

OWNER

Public reporting burden for this collection of information is estimated to average 5 hours per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send COMMENTS regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to:

Reports Clearance Officer, PHS Hubert H. Humphrey Building, Room 721-B 200 Independence Avenue, S.W. Washington, DC 20201

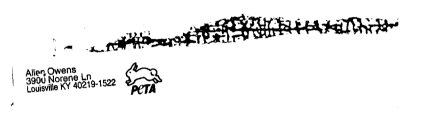
and to:

Office of Management and Budget Paperwork Reduction Project (0910-0025)

Washington, DC 20503

Attn: PRA

Please DO NOT RETURN this application to either of these addresses.





Dockets Management Branch (HFA-305)
Food & Drug Administration
Room # 63
5600 Fishers Lane
Rockville, MD 20857